

# Pine Chemicals Association, Inc.

March 15, 2002

Honorable Christine Todd Whitman Administrator US EPA P.O. Box 1473 Merrifield, VA 22116

Re: HPV Test Plan and Robust summaries for Fatty Acid Dimers and Trimer

Dear Ms Whitman;

On behalf or the member companies of the Pine Chemicals Association's High Production Volume Chemical Task Force, I am pleased to submit the Test Plan and Robust summaries for the following category;

"Farty Acid Dimers and Trimer"

The submission includes one electronic copy in pdf. Format, and a hard copy which will be mailed to EPA Headquarters. The Registration Number for our Consortium is

Should you have any questions concerning our submission please feel free to contact me at (770) 399-3112 or at wjones@pinechemicals.org.

Sincerely,

Walter L. Jones President & COO

> 1117 Perimeter Center West, Suite 6005 Atlanta, GA 30338

> > V: 770.399.3112 F: 770.399.3115 www.piechemicols.org

# 02 MAR 18 AM 11: 18 HIGH PRODUCTION VOLUME (HPV) CHEMICAL CHALLENGE PROGRAM

# **TEST PLAN**

for

# **FATTY ACID DIMERS AND TRIMER**

CAS No. 61788-89-4 CAS No. 68937-90-6 CAS No. 68783-41-5 CAS No. 71808-39-4

#### Submitted to the US EPA

By

The Pine Chemicals Association, Inc.
www.pinechemicals.org
HPV Task Force
Consortium Registration #

# **Table of Contents**

# **Test Plan for Fatty Acid Dimers and Trimer**

Sumn	nary	3
List of	FPCA HPV Consortium Members	6
i.	Description of Dimers and Trimer	7
	A. Composition  B. Commercial Uses  C. Complexity of Analytical Methodology	9
И.	Rationale for Selection of Representative Compound for Testing	11
III.	Review of Existing Data and Development of Test Plan	11
	<ul> <li>A. Evaluation of Physicochemical Data and Proposed Testing.</li> <li>B. Evaluation of Environmental Fate Data and Proposed Testing.</li> <li>C. Evaluation of Ecotoxicity Data and Proposed Testing.</li> <li>D. Evaluation of Human Health Effects Data and Proposed Testing.</li> </ul>	13 15
IV.	Robust Summaries of Existing Data	19

# Test Plan for Fatty Acid Dimers and Trimer

#### Summary

The Pine Chemicals Association, Inc. (PCA) is sponsoring 36 HPV chemicals. This Test Plan addresses the following four substances, known collectively as Fatty Acid Dimers and Trimer:

CAS Number	IUR Name	<b>Common Name</b>
61788-89-4	Fatty acids, C18-unsaturated, dimers	Dimer
68937-90-6	Fatty acids, C18-unsaturated, trimers	Trimer
68783-41-5	Fatty acids, C18-unsaturated, dimers, hydrogenated	Hydrogenated dimer
71808-39-4	Fatty acids, C16 and C18-unsaturated, dimerized	Crude dimer

All of the members of this category of substances (hereafter referred to by their common names throughout the test plan) are derived from unsaturated fatty acids, primarily tall oil fatty acids. As with other fatty acid-based products, these substances are complex mixtures and considered to be Class 2 substances.

Crude dimer is manufactured from C18 unsaturated fatty acids through heat treatment with or without an appropriate catalyst. The other members of this category are then obtained from crude dimer either by distillation and/or hydrogenation. (See **Figure 1** on page 8.)

The physical properties of all the members of this group are similar. They are all slightly viscous to viscous liquids and range in color from clear to dark brown. The largest end use for dimer is in the manufacture of polyamide resins for use in adhesives, inks and coatings.

PCA has reviewed existing data on these substances. There are existing data on dimer for many SIDS endpoints. These compounds are non-toxic in acute toxicity tests. Repeat-dose studies show low toxicity and no potential for reproductive effects. Bacterial and mammalian mutagenicity data are negative.

Where applicable, PCA will conduct physical/chemical property and environmental fate testing on all of the substances in the category for which data are not already available. PCA has elected to treat these four substances as a category for purposes of the HPV Program. Therefore, a representative of the category will be tested for the additional required SIDS endpoints. Dimer (CAS # 61788-89-4) has been selected as the representative substance in this category for testing as it has the largest production volume and, in particular the distilled form to be used for testing, has the highest dimer content of the members of this category.

A brief summary of the available data for the substances in this category, and the anticipated additional testing, is presented below in Table 1.

Table 1
Matrix of Available Adequate Data and Proposed Testing
On Fatty Acid Dimers and Trimer\*

	Required SIDS Endpoints										
Chemical and CAS#	Partition Coef.	Water Sol.	Biodeg.	Acute Fish	Acute Daph.	Acute Algae	Acute oral	Repeat Dose	In vitro genotox (bact.)	In vitro genotox (non- bact)	Repro/ Develop
Dimer 61788-89-4	Test	Test	Adeq.	Test	Test	Test	Adeq.	Adeq.	Adeq.	Adeq.	Adeq. Repro/ Test Develop.
Trimer 68937-90-6	Test	Test	Test	С	С	С	С	С	С	С	С
Hydrogenated dimer 68783-41-5	Test	Test	Test	С	С	С	С	С	С	С	С
Crude dimer 71808-39-4	Test	Test	Test	С	С	С	С	С	С	С	С

Adeq. Indicates adequate existing dataTest Indicates proposed testing

C Indicates category read-down from existing or proposed test data on dimer.

\* No testing will be conducted for melting point, boiling point, vapor pressure, hydrolysis, photodegradation and transport and distribution between environmental compartments as explained in the test plan.

# **Physical/Chemical Properties**

Physical and chemical properties will be determined for all members of the category when appropriate. However, many of the physical and chemical properties cannot be measured for these substances:

- Melting points will not be determined because these substances are complex mixtures and liquids under ambient conditions and have no specific melting point.
- Boiling points under ambient conditions cannot be determined because these substances are complex mixtures and will decompose before they boil.
- <u>Vapor pressure</u> of these substances under ambient conditions is essentially zero and experimental measurement is inappropriate.
- Water solubility of all of the substances in this category will be determined.

 <u>Partition coefficients</u> of all the substances in this category will be determined. However, the partition coefficient testing likely will yield more than one value representing the various components, rather than a single value representing the mixture.

#### **Environmental Fate**

With respect to the SIDS environmental fate endpoints:

- <u>Biodegradation</u> data will be generated for three of the substances in this category. Adequate data are already available for the fourth substance.
- Hydrolysis in water will not be determined for any of the substances in this category because the members of this category contain no hydrolyzable groups.
- <u>Photodegradation</u> is not relevant, since the vapor pressure of these substances is essentially zero and they could not enter the atmosphere.
- <u>Transport and distribution between environmental compartments</u> will not be determined due to the inability to provide usable inputs to the required model.

#### **Ecotoxicity**

 Using dimer, <u>acute toxicity to fish</u>, <u>daphnia and algae</u> will be tested under conditions that maximize solubility, but reduce exposure to insoluble fractions, which may cause nonspecific toxicological effects.

#### **Mammalian Toxicity**

- For the SIDS human health endpoints, there are adequate data on <u>acute</u> toxicity, repeat dose toxicity, reproductive effects and bacterial and <u>mammalian genotoxicity</u> for dimer.
- <u>Developmental toxicity</u> studies on dimer will be undertaken to fulfill this SIDS endpoint.

The Pine Chemicals Association, Inc. HPV Task Force includes the following companies:

Akzo Nobel Resins

Akzo Nobel - Eka Chemicals Incorporated

Arizona Chemical Company

Asphalt Emulsion Manufacturers Association

**Boise Cascade Corporation** 

Cognis Corporation

**Crompton Corporation** 

Eastman Chemical Co. (including the former Hercules Inc. Resins Division)

Georgia-Pacific Resins Inc.

Hercules Incorporated

ICI Americas (including the former Unigema)

Inland Paperboard & Packaging, Inc.

International Paper Co. (including the former Champion International

Corporation)

Koch Materials Co.

McConnaughay Technologies, Inc.

MeadWestvaco (including the former Mead Corp. and the former Westvaco)

Packaging Corporation of America

Plasmine Technology, Inc.

Raisio Chemicals

Rayonier

Riverwood International

Smurfit - Stone Container Corporation

Weyerhaeuser Co.

The Task Force has filed multiple test plans covering various chemicals. Not all members of the Task Force produce the substances covered by this test plan.

# I. Description of Fatty Acid Dimers and Trimer

The Pine Chemicals Association, Inc. (PCA) is sponsoring four HPV chemicals known collectively as Fatty Acid Dimers and Trimer. This category of chemicals consists of:

<b>CAS Number</b>	IUR Name	Common Name
61788-89-4	Fatty acids, C18-unsaturated, dimers	Dimer
68937-90-6	Fatty acids, C18-unsaturated, trimers	Trimer
68783-41-5	Fatty acids, C18-unsaturated, dimers, hydrogenated	Hydrogenated dimer
71808-39-4	Fatty acids, C16 and C18-unsaturated, dimerized	Crude dimer

For convenience, the common names of these substances are used in this test plan.

All the members of this category are produced by the dimerization of C18 unsaturated fatty acids, primarily tall oil fatty acids. As with other fatty acid-based products, these substances are complex mixtures and therefore are considered Class 2 substances.

The classical production process begins by heating unsaturated fatty acid in the presence of an acid-treated clay catalyst to a temperature greater than 200° C. Under these conditions, some of the fatty acids dimerize, a lesser amount trimerizes, and some isomerizes to monomer (Zinkel and Russell 1989). This reaction mixture is called crude dimer.

Based on EPA guidance (letter of May 4,1995), the reaction mixture can have two different CAS Registry Numbers depending on its intended use:

- If it is used as the feedstock for the production of dimer, it has CAS # 71808-39-4.
- If it is sold as a commercial raw material for the production of products other than dimer, then it has CAS # 61788-89-4.

The typical fatty acid dimerization process is shown in Figure 1 (see page 8).

MONOMER (Not part of this category) **HYDROGENATED** DIMER CRUDE DIMER 68783-41-5 **UNSATURATED** (REACTION **FATTY ACIDS** PRODUCT) 71808-39-4 Distillation Hydrogenation DIMER (standard) DISTILLED 61788-89-4 DIMER 61788-89-4 KNOWN AS DIMER WHEN SOLD COMMERCIALLY FOR OTHER PRODUCTS Distillation 61788-89-4 TRIMER 68937-90-6

Figure 1. Typical Fatty Acid Dimerization Process

Figure 1 shows schematically how the other members of this category of substances are produced from crude dimer.

- To produce the other members of the category, crude dimer is distilled, which
  removes most of the monomer, leaving dimer (known in the industry as
  standard dimer). The monomer acid is not a member of this category, but is
  included in PCA's Tall Oil Fatty Acid Test Plan.
- Standard dimer can be further distilled to give distilled dimer and trimer.
- Standard dimer and distilled dimer can also be hydrogenated to yield hydrogenated dimer, which has improved stability to heat and oxygen.

#### Composition

All the members of this category are liquids, ranging in color from clear to dark brown. The viscosity of the members depends on the dimer and trimer content, with crude dimer being a low viscosity liquid and at the other extreme, trimer being a very viscous liquid. The typical composition of the substances in this category is shown in Table 2 (below).

Table 2

Representative Compositions of Dimer Acids

Substance	C18 Acids (Monomeric) %	C36 Acids (Dimeric) %	C54 Acids (Trimeric) %	
Dimer				
Crude dimer Dimer Distilled dimer	30 2 1	60 80 94	10 18 5	
Trimer	<1	40	60	
Hydrogenated dimer	2	80	18	

The composition of dimer and trimer is complex. Several representative dimer structures are shown schematically in Figure 2 (see page 10). Dimers and trimers are predominantly cyclic addition compounds of unsaturated fatty acids, although bicyclic, non cyclic and other structures are present.

Figure 2. Representative Structures of Fatty Acid Dimers. (Many geometric isomers of the structures below are present in fatty acid dimers.)

# A. Commercial Uses of Dimer Acids

Dimer is used primarily in the production of non-nylon polyamide resins. Polyamide resins are used in various formulations for high performance adhesives and printing inks for flexible packaging, and as a cross linking agent for epoxy resins. Dimer is also used as a raw material for the production of surface coatings, primarily coatings for metal coils. Derivatives of dimer (imidazolines) are used as corrosion inhibitors in petroleum production and refining. Trimers find application as a component in oil well drilling fluids.

# B. Complexity of Analytical Methodology

All the substances in this category are Class 2 substances. This, combined with the fact that they are essentially insoluble in water and decompose rather than vaporize on heating at ambient pressure, creates a variety of analytical challenges. The large size of the dimer and trimer molecules and their non-volatility makes gas chromatography impractical. The commonly used technique for analyzing members of this category is gel permeation chromatography. However, this technique is not normally used for detecting low levels of dimer; rather, it is used for determining the monomer, dimer and trimer content of dimer products at percentage levels.

Preliminary method validation undertaken as a predicate to PCA's proposed HPV testing has indicated that under the correct conditions it should be possible to use this analytical method to determine levels of dimer as low as, or even <10 ppm, the expected solubility of these substances in water. While the level of dimer to be detected will be considerably higher in animal feeding studies, the analysis will be complicated by the presence of materials in the animal feed that are similar to dimer with respect to molecular weight and solubility. However, it is anticipated that this complication can be overcome and the selected technique will be satisfactory for the planned studies.

# II. Rationale for Selection of Representative Compound for Testing

Dimer (CAS # 61788-89-4) has been selected as the representative substance in this category for testing for the applicable SIDS ecotoxicity and developmental toxicity tests, as shown in Table 3 (identical to Table 1). Chemically, the members of this category are very similar as they are all essentially mixtures of monomer, dimer and trimer. Dimer has been selected as the representative test substance for ecotoxicity and developmental toxicity testing for several reasons. It is the most commercially important substance, with the largest production volume. Dimer, and in particular the distilled form to be used for testing, has the highest dimer content of any of the members of this category so that the results will be most representative of the category.

Since all of the substances in this category are either dimers or trimers of fatty acids, they are in the same family of compounds. Thus, these substances meet EPA's criterion of using the "family approach" to group chemicals into a category to examine related chemicals. In summary, these four substances -- all from the same family -- fit the requirements of the EPA's HPV Challenge Program for a category, and dimer is the most appropriate representative test material from this category.

# III. Review of Existing Data and Development of Test Plan

PCA has undertaken a comprehensive evaluation of all relevant data on the SIDS endpoints of concern for the substances in this category. Considerable data on dimer (CAS # 61788-89-4) are available that satisfy many of the SIDS endpoints for this category.

The availability of the data on the specific SIDS endpoints is summarized in Table 3 (identical to Table 1). Table 3 also shows data gaps that will be filled by additional testing and areas where data from dimer will be generalized to other category members.

Table 3
Matrix of Available Adequate Data and Proposed Testing
On Fatty Acid Dimers and Trimers\*

Observatoral	Required SIDS Endpoints										
Chemical and CAS #	Partition Coef.	Water Sol.	Biodeg.	Acute Fish	Acute Daph.	Acute Algae	Acute oral	Repeat Dose	In vitro genotox (bact.)	In vitro genotox (non- bact)	Repro/ Develop
Dimer 61788-89-4	Test	Test	Adeq.	Test	Test	Test	Adeq.	Adeq.	Adeq.	Adeq.	Adeq. Repro/ Test Develop.
Trimer 68937-90-6	Test	Test	Test	С	С	С	С	С	С	С	С
Hydrogenated dimer 68783-41-5	Test	Test	Test	С	С	С	С	С	С	С	С
Crude dimer 71808-39-4	Test	Test	Test	С	С	С	С	С	С	С	С

Adeq. Indicates adequate existing data

Test Indicates proposed testing

C Indicates category read-down from existing or proposed test data on dimer.

\* No testing will be conducted for melting point, boiling point, vapor pressure, hydrolysis, photodegradation and transport and distribution between environmental compartments as explained in the test plan.

# A. Evaluation of Existing Physicochemical Data and Proposed Testing

The basic physicochemical data required in the SIDS battery includes melting point, boiling point, vapor pressure, partition coefficient (K<sub>ow</sub>), and water solubility.

Such data are often meaningless for Class 2 substances such as those that comprise the dimer acid category. These substances are composed of a complex mixture of substances and are often difficult to characterize. The

members of this category are not only Class 2 substances, but are derived from natural sources. Their composition is variable and cannot be represented by a single chemical structural diagram. Due to this "complex mixture" characteristic of dimer, some physical property measurements, such as partition coefficient do not give single definitive results because the methodology used to determine these properties will actually fractionate or partition the substance into various components. Since the methodology will alter the actual sample composition, the results are likely to be erroneous, difficult to interpret, or meaningless.

# 1. Melting Point

Melting points will not be determined because all of the substances in this category are liquids under ambient conditions.

#### 2. Boiling Point

All of the substances in this category are produced at high temperatures, and generally by high vacuum distillation, and are non-volatile liquids at ambient temperatures. A boiling point at ambient temperature has no significance because these materials will thermally decompose before they boil. Accordingly, measurement of this property is inappropriate for all the substances in this category.

#### 3. Vapor Pressure

Vapor pressures for the substances in this category are effectively zero at ambient temperatures, and their experimental measurement is inappropriate.

#### 4. Water Solubility

Assuming adequate analytical sensitivity can be achieved, the water solubility of all of the substances in this category using OECD (105) will be determined.

#### 5. Partition Coefficient

The partition coefficient (i.e.,  $K_{ow}$ ) for all of the substances in this category will be determined using OECD method 107. It is likely that more than one  $K_{ow}$  value, rather than a single value, will be generated when this endpoint is determined. This outcome reflects the complex nature of Class 2 mixtures.

Summary of Physicochemical Properties Testing: The water solubility (OECD method 105) and partition coefficients (OECD method 107) of all of the substances in this category will be determined. Tests for melting point, boiling point, and vapor pressure are inapplicable to these substances.

# B. Evaluation of Existing Environmental Fate Data and Proposed Testing

The fate or behavior of a chemical in the environment is determined by the reaction rates for the most important transformation (degradation) processes. The basic environmental fate data covered by the HPV Program include biodegradation, stability in water (hydrolysis as a function of pH), photodegradation and transport and distribution between environmental compartments.

# 1. Biodegradation

Biodegradability provides a measure for the potential of compounds to be degraded by microorganisms. Depending on the nature of the test material, several standard test methods are available to assess potential biodegradability.

One of the chemicals in this category (dimer) has existing data on the biodegradation endpoint. Biodegradation for the other three substances will be determined using OECD protocol 301B.

#### 2. Hydrolysis

Hydrolysis as a function of pH is used to assess the stability of a substance in water. Hydrolysis is a reaction in which a water molecule (or hydroxide ion) substitutes for another atom or group of atoms present in an organic molecule. None of the substances in this category contain a functional group that would be susceptible to hydrolysis. Therefore, hydrolysis need not be measured.

In addition, low water solubility often limits the ability to determine hydrolysis as a function of pH. All of the substances in this category have very low solubility in water. Therefore, these materials are expected to be stable in water and it would be unnecessary to attempt to measure the products of hydrolysis.

#### 3. Photodegradation

Due to their lack of any vapor pressure under ambient conditions, there is essentially no opportunity for any of the fatty acid dimers or trimer to enter the atmosphere. Thus, photodegradation is irrelevant. In addition, based on the constituents in these complex mixtures, there is no reason to suspect that they would be subject to breakdown by a photodegradative mechanism. Consequently, this endpoint will not be determined for any of the substances in this category.

#### 4. Transport and Distribution between Environmental Compartments

The transport and distribution between environmental compartments is intended to determine the ability of a chemical to move or partition in the environment. The determination of this property requires the use of various models (e.g., level

III model from the Canadian Environment Modeling Centre at Trent University). For Class 2 substances such as dimer and related compounds, the required inputs to the model are either not available or not feasible to determine including molecular mass, reaction half-life estimates for air, water, soil, sediment, aerosols, suspended sediment, and aquatic biota. In addition, while the partition coefficient is also required and can be determined, the multiple  $K_{ow}$  values typically derived for these substances are a consequence of sample fractionation and reflect various components in the mixture and are not representative of the mixture itself. Consequently, due to the inability to provide usable inputs to the required model, no determination of transportation and distribution between environmental compartments will be undertaken for any of the substances in this category.

Summary of Environmental Fate Testing: Biodegradation data will be generated (using OECD 301B) for three of the compounds in this category; there are existing biodegradation data for dimer. Photodegradation, hydrolysis and transport and distribution between environmental compartments are not applicable to these substances.

#### C. Evaluation of Existing Ecotoxicity Data and Proposed Testing

The basic ecotoxicity data that are part of the HPV Program include acute toxicity to fish, daphnia and algae. Dimer will be tested for these endpoints under conditions that maximize the solubility under the specific test exposure conditions, but reduce exposure to insoluble fractions, which may cause nonspecific toxicological effects. In addition, the effect of both filtering, to further minimize nonspecific physical effects, and of reducing the pH to the lower end of the acceptable range for test organism survival, will also be investigated for changes in toxicological effects. The results of preliminary tests will be used to select the most appropriate test conditions for the definitive test for each species.

Summary of Ecotoxicity Testing: The acute toxicity of distilled dimer to fish (OECD 203), daphnia (OECD 202) and algae (OECD 201) will be tested under conditions that maximize solubility, but reduce exposure to insoluble fractions, which may cause nonspecific toxicological effects.

#### D. Evaluation of Existing Human Health Effects Data and Proposed Testing

#### 1. Acute Oral Toxicity

Acute oral toxicity studies investigate the effect(s) of a single exposure to a relatively high dose of a substance. This test is conducted by administering the test material to animals (typically rats or mice) in a single gavage dose. Harmonized EPA testing guidelines (August 1998) set the limit dose for acute oral toxicity studies at 2000 mg/kg body weight. If less than 50 percent mortality is observed at the limit dose, no further testing is needed. A test substance that

shows no effects at the limit dose is considered essentially nontoxic. If compound-related mortality is observed, then further testing may be necessary.

#### **Summary of Available Acute Oral Toxicity Data**

Dimer is non-toxic following acute oral exposure, with  $LD_{50}$  values > 2,000 mg/kg in several studies. Hydrogenated dimer is also non-toxic following acute oral exposure with an  $LD_{50}$  value > 5,000 mg/kg.

Summary of Acute Oral Toxicity Testing: The representative compound in this category (dimer) has been tested for acute oral toxicity and found to be non-toxic (i.e.,  $LD_{50} > 2000 \text{ mg/kg}$ ). In addition, hydrogenated dimer is also non-toxic with an  $LD_{50}$  value > 5,000 mg/kg.

#### 2. Repeat Dose Toxicity

Subchronic repeat dose toxicity studies are designed to evaluate the effect of repeated exposure to a chemical over a significant period of the life span of an animal. Typically, the exposure regimen in a subchronic study involves daily exposure (at least 5 consecutive days per week) for a period of not less than 28 days or up to 90 days (i.e., 4 to 13 weeks). The HPV program calls for a repeat dose test of at least 28 days. The dose levels evaluated are lower than the relatively high doses used in acute toxicity (i.e., LD<sub>50</sub>) studies. In general, repeat dose studies are designed to assess systemic toxicity, but the study protocol can be modified to incorporate evaluation of potential adverse reproductive and/or developmental effects.

#### **Summary of Available Repeat Dose Toxicity Data**

There are existing data that demonstrate low toxicity for dimer. This substance was administered to Sprague-Dawley rats at dietary concentrations of 0, 0.1, 1, or 5% for 13 weeks. The approximate doses were 0, 100, 1,000, or 5,000 mg/kg/day. Parameters evaluated included clinical signs, body weight, food and water consumption, hematology, clinical chemistry, and gross pathology, organ weights and microscopic pathology.

No deaths occurred and no treatment-related effects on clinical signs, body weight, body weight gain, or water intake were noted. A transient, statistically significantly decrease in food consumption occurred in the 5% males and females during the first four weeks of study. Slight changes in hemoglobin (increased in 5% males) and prothrombin time (increased in 1% females and 5% males and females) were considered not to be toxicologically significant. Treatment-related clinical chemistry changes included increased alkaline phosphatase (1 and 5% males and females) and ALT (5% males and females), and decreases in total cholesterol and triglycerides (1 and 5% males and females), total serum protein and albumin (5% males and females), and beta-

globulin fraction (1 and 5% males). While some decreased organ weights were noted, they did not correlate to any microscopic changes. Although a no-effect-level was not identified in this study, 0.1% (approximately 100 mg/kg/day) can be considered a no-observed-adverse-effect-level (NOAEL) based on minimal increases in clinical chemistry parameters and histopathological findings at the higher doses.

Summary of Repeat Dose Toxicity Testing: Dimer has been tested for repeat dose toxicity in a 13 week study. In this study, the NOAEL was approximately 100 mg/kg/day, indicating that this compound has low toxicity.

#### 3. Genotoxicity - In vitro

Genetic testing is conducted to determine the effects of substances on genetic material (i.e., DNA and chromosomes). The gene, which is composed of DNA, is the simplest functional genetic unit. Mutations of genes can occur spontaneously or as a consequence of exposure to chemicals or radiation. Genetic mutations are commonly measured in bacterial and mammalian cells, and the HPV program calls for completing both types of tests.

#### **Summary of Available Genotoxicity Data**

Dimer has been tested for potential genotoxicity in several test systems including the Ames *Salmonella* assay, mouse lymphoma cell assay and a metaphase chromosome analysis of human lymphocytes. None of these test systems showed any indication of genotoxicity.

Summary of Genotoxicity Testing: Because dimer has been tested and found negative in three genotoxicity assays, no additional testing for this endpoint will be undertaken.

# 4. Reproductive and Developmental Toxicity

Reproductive toxicity includes any adverse effect on fertility and reproduction, including effects on gonadal function, mating behavior, conception, and parturition. Developmental toxicity is any adverse effect induced during the period of fetal development, including structural abnormalities, altered growth and post-partum development of the offspring.

The "toxicity to reproduction" aspect of the HPV Challenge Program can be met by conducting a reproductive/developmental toxicity screening test or adding a reproductive/developmental toxicity screening test to the repeat dose study (OECD 421 or OECD 422, respectively).

#### Summary of Reproductive/Developmental Toxicity Data

As noted in the SIDS guidelines for the reproduction toxicity endpoint, "when a 90-day repeated dose study is available and demonstrates no effects on the reproductive organs, in particular the testes, then a developmental study can be considered as an adequate test to complete information on reproduction/developmental effect." Dimer has been tested in a 13-week repeat dose study. This study included histopathology of reproductive organs (i.e., testes, ovaries, uterus) and showed no evidence of reproductive organ toxicity at any dose level. Therefore, this study satisfies the SIDS reproductive toxicity endpoint.

Summary of Reproductive/Developmental Testing: Dimer did not demonstrate any effects on reproductive toxicity in a repeat dose study. However, since this study did not evaluate potential developmental toxicity, dimer will be tested for this endpoint with OECD method 421.

# References

Zinkel, D.F. and Russell, J., Eds. 1989. Naval Stores. Production, Chemistry, Utilization. Pulp Chemicals Association, New York.

March 2002



# Robust Summaries of Existing Data 18 AHII: 19

Test Substance	
Chemical Name	Fatty acids, C18-unsaturated dimers
CAS#	61788-89-4
· · · · · · · · · · · · · · · · · · ·	County programming the contract of the contrac
Remarks	This substance is referred to as dimer in the test plan for Fatty Acid Dimers and Trimer
<u>Method</u>	
Method/Guideline followed	Testing was conducted according to OECD Test Method 301B "Ready Biodegradability: Modified Sturm Test."
Test Type	Aerobic
(aerobic/anaerobic)	
GLP (Y/N)	Y
Year (Study Performed)	1991
Contact time	28 days
Inoculum	Activated sludge from a municipal sewage treatment plant
	Promatod oldago from a marifolipal sewage treatment plant
Test conditions	Inoculum: Activated sludge microorganisms were obtained from a municipal sewage treatment plant at Wateschap de Aa, Schijndel, the Netherlands.
	Concentration of test chemical: The test material was used at concentrations of 10 and 20 mg/L.
	Test Setup: Nutrient medium was prepared by adding 2 mL of a potassium phosphate solution, 1 mL each of magnesium sulfate, calcium chloride, and ammonium sulfate solutions, and 4 mL of a ferric chloride solution to a final volume of 1 liter in purified water. Twenty-four hours prior to study initiation, vessels were filled with the nutrient culture medium and 30 mL of inoculum and aerated over night. On day 1 of the study, the test and reference material (sodium benzoate, 20 mg/L) were added and the total volume was increased to 3 liters. The following series of vessels was prepared: culture medium with inoculum; culture medium with inoculum and sodium benzoate; and culture medium with inoculum and test material. The CO <sub>2</sub> absorption bottles were connected in series to the exit air line of each bottle. CO <sub>2</sub> -free air was bubbled through the solution. All experiments were performed at 20 to 22°C.
	Sampling frequency: Samples were collected from the first CO <sub>2</sub> absorber vessel on days 2, 5, 7, 9, 12, 16, 21, and 28.
	Controls: Yes.
	Analysis: Samples from the CO <sub>2</sub> absorbers were analyzed using a Heraeus CHN-analyzer.

Results	
Degradation % after time	6.6% at 10 mg/L and 6.3% at 20 mg/L at 28 days (test article); 71% at 28 days (sodium benzoate).
Conclusions	The test article, at low and high concentrations, was degraded approximately 6% after 28 days and sodium benzoate was degraded 71% after 28 days. Under the conditions of the OECD guidelines, the test article was not readily biodegradable.
Data Quality	Reliable without restrictions— Klimisch Code 1a
<u>Reference</u>	Coenen, T.M.M. 1991. Ready biodegradability: modified Sturm test. RCC NOTOX Project 052559. NOTOX, The Netherlands.

ACUTE TOXICITY – ORAL				
Test substance				
Chemical Name CAS #	Fatty acids, C18-unsaturated dimers			
Remarks	This substance is referred to as dimer in the test plan			
	for Fatty Acid Dimers and Trimer			
<u>Method</u>				
Method/Guideline followed	Testing was conducted according to OECD Test Method 401, "Acute Oral Toxicity."			
GLP (Y/N)	Υ			
Year (Study Performed)	1986			
Species	Rat			
Strain	Wistar			
Route of administration	Oral			
Dose levels	5,000 mg/kg			
Sex and number/group	5 male and 5 female rats			
Frequency of treatment	Single oral gavage			
Duration of test	14 day observation post-treatment			
Control group (Y/N)	N			
Result				
Acute Oral LD <sub>50</sub>	>5,000 mg/kg			
<u>Detailed Summary</u>	Wistar rats (n = 5/sex) received a single oral dose of 5000 mg/kg of dimer (CAS #61788-89-4) and were observed for 14 days. Parameters evaluated included clinical signs, mortality, body weight, and gross pathology. No effects on mortality, clinical signs or body weight were reported. Gross necropsy revealed no treatment-related effects. The acute oral LD <sub>50</sub> was greater than 5000 mg/kg.			
Data Quality	Valid without restriction – Klimisch Code 1a			
<u>Reference</u>	Thouin, M.H. 1986. Evaluation of acute oral toxicity of			
	[trade name deleted; dimer] in the rat. NOTOX 0336/416.			
	NOTOX, The Netherlands.			

ACUTE TOXICITY - ORAL	
Test substance	
Chemical Name	Fatty acids, C18-unsaturated dimers
CAS#	61788-89-4
Remarks	This substance is referred to as dimer in the test plan
	for Fatty Acid Dimers and Trimer
<u>Method</u>	
Method/Guideline followed	Testing was conducted according to OECD Test Method
	401, "Acute Oral Toxicity."
GLP (Y/N)	Υ
Year (Study Performed)	1989
Species	Rat
Strain	Sprague-Dawley
Route of administration	
Dose levels	2,000 mg/kg
Sex and number/group	5 male and 5 female rats
Frequency of treatment	Single oral gavage
Duration of test	14 day observation post-treatment
Control group (Y/N)	N
Result	> 0.000 ···· · #··
Acute Oral LD <sub>50</sub>	>2,000 mg/kg
<u>Detailed Summary</u>	Sprague-Dawley rats (n = 5/sex) received a single oral
	dose of 2000 mg/kg of dimer (CAS #61788-89-4) and were
	observed for 14 days. Parameters evaluated included
	clinical signs, mortality, body weight, and gross pathology.
	No mortalities occurred and no changes in clinical signs,
	body weight or gross pathology were reported. The acute
Data Quality	oral LD <sub>50</sub> was greater than 2000 mg/kg. Valid without restriction – Klimisch Code 1a
Reference	
Velerelice	Saboureau, D. 1989. Evaluation of the acute toxicity in the
	rat by the oral route. TAO 88.1518. Biogir S.A. Conseil Recherche, France.
	recherche, mance.

ACUTE TOXICITY - ORAL	
Test substance	
Chemical Name CAS #	Fatty acids, C18-unsaturated dimers, hydrogenated 68783-41-5
Remarks	This substance is referred to as hydrogenated dimer in the test plan for Fatty Acid Dimers and Trimer
Method	
Method/Guideline followed	Test procedure was consistent with OECD Test Method 401, "Acute Oral Toxicity."
GLP (Y/N)	Υ
Year (Study Performed)	1988
Species	Rat
Strain	Wistar
Route of administration	Oral
Dose levels	5,000 mg/kg
Sex and number/group	5 male and 5 female rats
Frequency of treatment	Single oral gavage
Duration of test	14 day observation post-treatment
Control group (Y/N)	N
Result	
Acute Oral LD <sub>50</sub>	>5,000 mg/kg
<u>Detailed Summary</u>	Wistar rats (n = $5/\text{sex}$ ) received a single oral dose of 5000 mg/kg of hydrogenated dimer (CAS #68783-41-5) and were observed for 14 days. Parameters evaluated included clinical signs, mortality, body weight, and gross pathology. Mortality, clinical signs, body weight, and gross pathology were unaffected by treatment. The acute oral LD <sub>50</sub> was greater than 5000 mg/kg.
Data Quality	Valid without restriction – Klimisch Code 1a
<u>Reference</u>	Reijnders, J.B.J. 1988. Acute oral toxicity of [trade name deleted; hydrogenated dimers] in the rat. RCC NOTOX 0811/1041. NOTOX, The Netherlands.

REPEAT DOSE TOXICITY	
Test substance	
Chemical Name	Fatty acids, C18-unsaturated dimers
CAS#	61788-89-4
Remarks	This substance is referred to as dimer in the test plan
	for Fatty Acid Dimers and Trimer
<u>Method</u>	
Method/Guideline followed	Testing was conducted according to OECD Test Method 408, "Subchronic Oral Toxicity – Rodent: 90-Day."
Year	1993
GLP (Y/N)	Y
Species	Rat
Strain	Sprague-Dawley
Sex	Male, female
Route of Administration	Oral, diet
Exposure Period	13 weeks
Frequency of Treatment	Daily
Post-exposure observation	None
period	
Dose Levels	0.1, 1, 5%
Control group (Y/N)	According to the Commence of t
Results	
NOAEL:	0.1%
Detailed Summary	Dimer (CAS #61788-89-4) was administered to CD Sprague-Dawley rats (n = 20/sex/group) in the diet at concentrations of 0, 0.1, 1, or 5% for 13 weeks. The approximate doses were 0, 100, 1,000, or 5,000 mg/kg/day, based on standard conversion factors (WHO 1990). Parameters evaluated included clinical signs, body weight, food and water consumption, ophthalmoscopy, hematology, clinical chemistry, gross pathology, organ weights (brain, heart, liver, kidneys, spleen, testes, adrenal glands), and microscopic pathology (adrenal glands, brain, colon, femur and stifle joint, ileum, larynx, lymph nodes, muscle, ovaries and fallopian tubes, pituitary, sciatic nerve, sternum, thyroid and parathyroids, uterus, aorta, cecum, duodenum, head, jejunum, liver, esophagus, pancreas, prostate, spinal cord, stomach, tongue, bladder, cervix, heart, kidneys, lungs, mammary glands, rectum, spleen, thumus, trachea, epididymides, skin, salivary glands, testes, seminal vesicles, vagina, eyes/harderian glands).  No deaths occurred and no treatment-related effects on clinical signs, body weight, body weight gain, water intake, or ophthalmoscopy were noted. A transient, statistically significantly decrease in food consumption occurred in the 5% males and females during the first four weeks of study. The animals exhibited normal consumption from week 4 through 13. Slight changes in hemoglobin (increased in 5%

	males) and prothrombin time (increased in 1% females and
	5% males and females) were considered not to be
	toxicologically significant. Treatment-related clinical
	chemistry changes included statistically significant
	increases in alkaline phosphatase (1 and 5% males and
	females) and ALT (5% males and females), and statistically
	significant decreases in total cholesterol (1 and 5% males
	and females), triglycerides (1% males and 5% males and
	females), total serum protein and albumin (5% males and
	females), and beta-globulin fraction (1 and 5% males). At
	necropsy, the mesenteric lymph nodes were slightly to
	moderately enlarged in all dimer treatment groups and the
	incidence of uterine fluid distension was increased at 5%.
	Absolute and relative spleen (males at 1 and 5%) and liver
	(males and/or females at 1 and 5%) weights were
	statistically significantly decreased. In addition, absolute
	kidney weight was significantly decreased in females at 5%
	and absolute and relative liver weights were significantly
	decreased in females at 0.1%. The relevance of these
	decreases in organ weights is not known, since they did not
	correlate to any microscopic changes. Histopathology
	revealed treatment-related findings in the following organs:
	mesenteric lymph nodes (aggregation of macrophages in
	both sexes at 0.1% and higher); spleen (macrophages with
	brown pigment in both sexes at 1 and 5% and in the
	females at 0.1%); liver (bile duct proliferation and bile duct
	sclerosis in males at 5%); adrenals (cortical vacuolation in
	females at 1 and 5%); and thyroids (follicular epithelial
	hypertrophy in females at 5%).
	Although a no-effect-level was not identified in this study,
	0.1% (approximately 100 mg/kg/day) can be considered a
	no-observed-adverse-effect-level based on increases in
	clinical chemistry parameters and histopathological findings
	at the higher doses.
	Valid without restriction – Klimisch Code 1a
	Spurgeon, M., and Hepburn, P. 1993. Dimer acid: 13
	week feed study in the rats. Study FT920485.
ĺ	Environmental Safety Laboratory, England.
l	World Health Organization (WHO). 1990. Principles for
	the Toxicological Assessment of Pesticide Residues in
	Food.

Data Quality References

IN VITRO GENETIC TOXICITY			
<u>Test substance</u>			
Chemical Name	Fatty acids, C18-unsaturated dimers		
CAS#	61788-89-4		
Remarks	This substance is referred to as dimer in the test plan for Fatty Acid Dimers and Trimer		
<u>Method</u>			
Method/Guideline followed	Testing was conducted according to OECD method # 471, "Bacterial Reverse Mutation Assay"		
Year	2000		
GLP (Y/N)	Yes		
System of testing	S. typhimurium strains TA98, TA100, TA 102, TA1535, TA1537		
Concentrations	33, 100, 333, 1000, 2500 and 5000 μg/plate		
Metabolic activation	With and without S9		
Results	Non-mutagenic		
<u>Detailed Summary</u>	An Ames test was conducted in <i>S. typhimurium</i> strains TA98, TA100, TA 102, TA1535, and TA1537. Dimer (CAS #61788-89-4) concentrations of 33, 100, 333, 1000, 2500 and 5000 µg/plate were tested with and without metabolic activation (S9 mix). No substantial increase in revertant colony numbers of any of the five tester strains was observed following treatment with dimer at any concentration level either in the presence or absence of metabolic activation (S9 mix). Thus, dimer was not considered to be mutagenic.		
Data Quality	Valid without restriction – Klimisch Code 1a		
Reference	Wollny, H. 2000. Salmonella Typhimurium assay with [trade name deleted; dimers] RCC Cytotest Cell Research, GMBH, Robdorf.		

IN VITRO GENETIC TOXICITY	1
Test substance	
Chemical Name	Fatty acids, C18-unsaturated dimers
CAS#	61788-89-4
Remarks	This substance is referred to as dimer in the test plan
	for Fatty Acid Dimers and Trimer
<u>Method</u>	
Method/Guideline followed	Testing was conducted according to OECD Test Method
V	476, "In vitro Mammalian Cell Gene Test."
Year	Y
GLP (Y/N)	200 - 200 -
System of testing	Mouse lymphoma L5178Y cells
Concentration	25 to 300 μg/mL
Metabolic activation	With and without
Results	Non-mutagenic
<u>Detailed Summary</u>	Dimer (CAS #61788-89-4) was incubated in vitro with
	L5178Y mouse lymphoma cells for three hours at
	concentrations ranging from 25 to 300 μg/mL with and
	without metabolic activation (S9 mix). Samples were
	collected at 24 and 48 hours to assess growth. After 48
	hours, cells were collected, plated, and incubated for 12
	days to assess viability and mutant frequency. The assay
	was conducted in duplicate.
	In Test 1 (without S9), toxicity was observed at 300 $\mu$ g/mL and in Test 2 (without S9) toxicity was observed at 275 and 300 $\mu$ g/mL. These concentrations were excluded from the mutation analyses. A statistically significant increase in mutant frequency was observed in Test 2 at 250 $\mu$ g/mL. However, because the increase was small, it was not considered biologically significant; no increase occurred in Test 1. Tests 1 and 2 (with S9 mix) produced reduced survival at 300 $\mu$ g/mL and 250 $\mu$ g/mL and above, respectively. These concentrations were excluded from the mutation analyses. No increase in mutant frequency was observed. Dimer acid did not demonstrate mutagenic potential in this assay.
Data Quality	Valid without restriction – Klimisch Code 1a
Reference	Adams, K. 1993. Dimer acid: mouse lymphoma TK locus
	assay. ULR 472/930202. Huntingdon Research Centre
	Ltd., England.

IN VITRO GENETIC TOXICITY			
Test substance			
Chemical Name	Fatty acids, C18-unsaturated dimers		
CAS#	61788-89-4		
Remarks	This substance is referred to as dimer in the test plan		
	for Fatty Acid Dimers and Trimer		
<u>Method</u>			
Method/Guideline followed	Testing was conducted according to OECD Test Method		
	473, "In Vitro Mammalian Cytogenetic Test."		
Year	1993		
GLP (Y/N)	Y		
System of testing	Human lymphocytes		
Concentration	9.4 to 300 μg/mL		
Metabolic activation	With and without		
<u>Results</u>	Non-mutagenic		
Detailed Summary	Human lymphocytes were incubated with dimer (CAS #61788-89-4) at concentrations ranging from 75 to 300 μg/mL with and without metabolic activation. In the first assay, the cultures containing S9 mix were centrifuged three hours after dosing and fresh medium was added for an additional 15 hours. In the second assay, half the cultures were processed following the procedure used in the first assay with a harvest at 18 hours and the other half were harvested at 32 hours. For all tests, two hours prior to treatment cessation, mitotic activity was arrested by the addition of colchicine, and the number of mitotic cells per 1000 cells in each culture was determined microscopically. No significant increase in the proportion of aberrant cells was observed in either the first or second assay with or without metabolic activation. Dimer demonstrated no clastogenic activity in this assay.		
Data Quality	Valid without restriction – Klimisch Code 1a		
Reference	Akhurst, L. 1993. Dimer acid: metaphase chromosome		
	analysis of human lymphocytes cultured <i>in vitro</i> . ULR		
	471/930241. Huntingdon Research Centre Ltd., England.		